(12) INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(19) World Intellectual Property Organization International Bureau



(43) International Publication Date 21 August 2003 (21.08.2003)

(10) International Publication Number WO 03/068968 A2

- (51) International Patent Classification7: C12N 15/12, 15/63, C07K 14/47, A01K 67/027, C07K 16/18, G01N 33/68
- (21) International Application Number: PCT/IB03/01044
- (22) International Filing Date: 14 February 2003 (14.02.2003)
- (25) Filing Language:

English

(26) Publication Language:

English

(30) Priority Data:

60/356,136

14 February 2002 (14.02.2002) US

02290610.1 11 March 2002 (11.03.2002) EP

- (71) Applicant (for all designated States except US): IN-STITUT NATIONAL DE LA SANTE ET DE LA RECHERCHE MEDICALE (INSERM) [FR/FR]; 101, rue de Tolbiac, F-75654 Paris Cédex 13 (FR).
- (72) Inventors; and
- (75) Inventors/Applicants (for US only): MALISSEN, Marie [FR/FR]; 11 avenue de la Pinede, 13009 Marseille (FR). MALISSEN, Bernard [FR/FR]; 11 avenue de la Pinede, 13009 Marseille (FR). AGUADO VIDAL, Enrique [ES/ES]; c/o Alonso de Ojeda, No., 4C, 30007 Murcia (ES).

- (74) Agent: BECKER, Philippe; Cabinet BECKER ET AS-SOCIES, 35 rue des Mathurins, F-75008 Paris (FR).
- (81) Designated States (national): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW.
- (84) Designated States (regional): ARIPO patent (GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, SE, SI, SK, TR), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).

Declaration under Rule 4.17:

of inventorship (Rule 4.17(iv)) for US only

Published:

without international search report and to be republished upon receipt of that report

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

(54) Title: MUTATED GENE CODING FOR A LAT PROTEIN AND THE BIOLOGICAL APPLICATIONS THEREOF

(57) Abstract: The present invention relates to a mutated gene coding for a mutant LAT protein leading to an exaggerated T_H2 differentiation. The invention relates to biological structures containing said mutant, particularly, non-human LAT gene mutated animals, plasmids, chromosomal DNA's, embryos comprising said mutated gene, and applications thereof. The invention further relates to screening method for drug useful for treatment against asthma and allergy. Otherwise, the invention relates to method for producing IgE antibodies.



Mutated gene coding for a LAT protein and the biological applications thereof.

The present invention relates to a mutated gene coding for two mutant LAT proteins leading to an exaggerated $T_{\rm H}2$ cell differentiation. The invention relates to biological structures containing said mutant, particularly, non-human 5 LAT gene mutated animals, cell cultures, plasmids, chromosomal DNAs, embryos comprising said mutated gene, and applications thereof. The invention further relates to screening methods for drug useful for treatment against asthma, allergy and any pathological immune responses involving $T_{\rm H}2$ cells. The invention also relates to method for producing IgE antibodies.

Background Art

A key event in the pathogenesis is the production of antibodies of the IgE class. Hypergammaglobulinemia E results from loss of immunoregulation. More specifically, T lymphocyte abnormalities have been reported in a number of pathologic hyper IgE conditions and are the object of much research aiming at developing pharmaceutical compounds that 20 will prevent atopic allergy and asthma.

TCR recognize peptide fragments bound major histocompatibility complex (MHC) molecules and relay this information to the interior of the T cell via adapter 25 proteins. One of these, the adapter LAT (Linker for Activation of T cells), coordinates the assembly of signaling complexes through multiple tyrosine residues intracytoplasmic segment. Upon within its TCR-induced phosphorylation, each of these tyrosine residues manifests 30 some specialization in the signaling proteins it recruits. Studies on cell lines showed that mutation of tyrosine 136 (Y136) selectively eliminates binding of phospholipase Cyl (PLC-y1) whereas the simultaneous mutation of Y175, Y195 and

Y235 results in loss of binding of downstream adapters Gads and Grb-2 (Lin and Weiss, 2001; Samelson et al, 1999; Zhang et al, 2000). Studies of LAT "knock in" mutant mice presenting the mutation of the four distal tyrosine residues of LAT in phenylalanine, called 4YF mice, showed that the murine T cell development was completely blocked (Sommers et al,2001). Hence, their thymocyte development was arrested at the immature CD4 CD8 stage and no mature T cells were present.

10

The present invention now provides genetic evidence that LAT exerts an unanticipated and surprising inhibitory function on the differentiation of CD4 helper T (T_H) cells into T_H2 cells.

15

Mice homozygous for the mutation of a single LAT tyrosine (LAT Y136F) results in mice that show a precocious and spontaneous accumulation of polyclonal $T_{\rm H}2$ cells, which chronically produce large amounts of interleukins 4, 5, 10 and 13. This exaggerated $T_{\rm H}2$ differentiation leads in turn to tissue eosinophilia and to the maturation of massive numbers of plasma cells secreting IgE and IgG1 antibodies (see Figure 1). Thus, in addition to known positive signaling, LAT also appears essential for establishing inhibitory signals that control T cell homeostasis.

Mice for the composite mutation of the three distal LAT tyrosines (LAT Y175F+Y195F+Y235F) prevents the development of T cells expressing alpha/beta T cell receptor. However, it allows the development of T cells expressing gamma/delta T cell receptors, and their accumulation in the periphery (see figure 9). These polyclonal gamma/delta T cells chronically produce large amounts of interleukins 4, 5, 10 and 13 (i.e. they present blatant TH2 phenotype). This exaggerated TH2-type differentiation of gamma/delta T cells leads in turn to the maturation of massive numbers of plasma cells secreting IgE and IgG1 antibodies (see Figures 10 and

11).

Description of drawings

Figure 1 is a diagram disclosing the immune system 5 development of mutant mice.

Figure 2 illustrates the LAT Y136F knock-in strategy:

- (1) : the partial restriction map of the wild-type LAT gene.
- (2): the targeting vector used for the introduction of 10 the LAT Y136F mutation.
 - (3): the structure of the targeted allele following homologous recombination.
 - (4) : the final structure of the targeted allele after removal of the ${\rm neo^r}$ gene via Cre-mediated recombination.
- 15 Figure 3 illustrates the aberrant growth of lymphoid organs in the mice: thymus (A), spleen (B) and lymph nodes (C).

 Figure 4 relates to constitutive type-2 cytokine production in CD4 T cells freshly isolated from LAT peripheral lymphoid organs.
- 20 Figure 5 relates to a phenotypic analysis of T cells from wild-type and LAT^{Y136F} mice.
 - Figure 6 illustrates eosinophilia in 6 weeks old LATY136F lymphoid organs.
- A: Dot plot panel showing the gate selected for the analysis described in panel B and for the sorted cells picture in panel C.
 - B: Single color histograms of gated cells labelled with antibodies characterizing eosinophils.
 - C: Hematoxylin and eosin staining of sorted cells.
- 30 Figure 7 illustrates the hyperactivity of B lymphocytes: massive serum levels of IgE and IgG1 antibodies in unimmunized LAT^{Y136F} mice.
 - Figure 8 illustrates the LAT Y175F+Y195F+Y235F knock in strategy:
- 35 (1): the partial restriction map of the wild-type LAT gene.
 - (2) : the targeting vector used for the introduction of



the LAT Y175F, Y195F and Y235F mutation.

- (3): the structure of the targeted allele following homologous recombination.
- (4): the final structure of the targeted allele after 5 removal of the neor gene via Cre-mediated recombination. Figure 9 relates to a phenotypic analysis of the gamma/delta T cells developed in large numbers in the LAT Y175F+Y195F+Y235F mutant in the mere absence of alpha/beta T cells.
- 10 Figure 10 illustrates the $T_{\rm H}2$ -type cytokines that are spontaneously produced by the gamma/delta T cells present in LAT Y175F+Y195F+Y235F mutant mice (lane 3) and compare them the $T_{\rm H}2$ -type cytokines that are spontaneously produced by the alpha/beta T cells developed in the LAT Y136F mutant (lane 15 1).

Figure 11 illustrates the hyperactivity of B lymphocytes and the massive amounts of IgE anf IgG1 that are spontaneously found in the serum of unimmunized LAT Y175F+Y195F+Y235F mice.

20

Description

In this application, LAT Y136F, LAT Y175F, LAT Y195F, and LAT Y235F refer to the designated mutation itself, while LAT Y136F, LAT Y175F, LAT Y195F and LAT Y235F refer to mutants, mice 25 or products derived from these mutations.

Mutation of one or three tyrosine(s) among the four distal tyrosine of LATprotein (i.e. LAT Y136F, or LAT Y175F+Y195F+Y235F) is able to induce the development of associated with exacerbated $T_{H}2$ immunity. 30 pathologies Characteristics of the phenotype associated with this mutation are described in the following examples. Therefore, the present invention provides models of allergy and/or associated with or other diseases $T_{\rm H}2$ cell asthma 35 deregulation or activity, more particularly $T_{\rm H}2$ cell accumulation. Among the advantages of said models, it is found the rapidity of the model preparation (about 3-4 weeks

for a mice model instead of several months) and the exacerbated phenotype (for instance, exacerbated IgE production and tissue eosinophilia).

5 This phenotype due to the mutation of one or three tyrosine(s) among the four distal tyrosine residues of LAT protein (namely, LAT Y136F, or LAT Y175F+Y195F+Y235F) in mice was unpredictable, considering the phenotype of mice in which the four mutations are combined (LAT 4YF mice). 10 Indeed, LAT 4YF mice are totally devoid of thymocytes and T cells, because of the early differentiation blockage. Therefore, the LAT 4YF mice are unable to lead or suggest the phenotype observed for the LAT^{Y136F} , and $LAT^{Y175F+Y195F+Y235F}$ mice. Moreover, none of the results of the previous studies 15 on cell lines suggests such a phenotype. Furthermore, the phenotype obtained in mice with the mutation Y136F could not be extrapolated in order to deduce the expected phenotype of mice having a composite mutation Y175F+Y195F+Y235F because of the different effects of the mutation Y136F and the 20 mutations Y175F, Y195F, and Y235F observed during the cell line studies.

The object of the present invention is to provide non-human animals having a mutated LAT gene of the invention leading to an exaggerated TH2 cell differentiation. By "gene" is intended cDNA or genomic sequence coding for a LAT protein. By "mutated LAT gene of the invention" is intended a LAT gene coding for a mutant LAT protein, the sequence of which corresponds to a wild type sequence and contains the mutation of a single tyrosine among the four distal ones corresponding to Y136 in the mouse LAT protein or the composite mutation of the three distal tyrosine residues (corresponding to Y175, Y195 and Y235 in the mouse LAT protein). For example, the tyrosine corresponding to Y136 in the mouse LAT protein. In a first preferred embodiment, said LAT gene coding for a mutant LAT protein contains a single mutation



of the tyrosine residue corresponging to Y136 in the mouse LAT protein. In a second preferred embodiment, said LAT gene coding for a mutant LAT protein contains the composite mutation of the three distal tyrosines, those corresponding 5 to Y175, Y195 and Y235 in the mouse LAT protein. Preferably, said non-human animals are mice and said non-human animals have the mutated gene coding for a mutant LAT protein, the sequence of which corresponds to a wild type sequence and contains the single mutation of the tyrosine residue at 10 position 136 or the composite mutation of the three distal tyrosine residues at positions 175, 195 and 235. Preferably, said mutation consists in the replacement by a residue preventing the association of the "tyrosine-based" sequences with the SH2 domain of proteins. More preferably, said 15 mutation consists in the replacement of the tyrosine by a phenylalanine (Y-F), an aspartic acid (Y-D) or a glutamic acid (Y-E). Still more preferably, said mutation consists in the replacement of the tyrosine by a phenylalanine (Y-F). Preferably, said non-human animals according 20 invention are mammals, and in particular, they are rodents. More preferably, said rodents are mice. Preferably, said animals are homozygous for the mutated LAT gene or are carrying a null allele of the LAT gene. Preferably, said mutated LAT gene is incorporated into the animal genome by 25 targeted insertion in order to keep said mutated LAT gene under the control of regulatory regions of the endogenous LAT gene.

By "distal" is intended the C-terminal end of the protein. 30 Therefore, the distal tyrosine residues are the tyrosines residues located at the C-terminal end of the protein.

In particular, the invention concerns any germ cell and somatic cell from said animals or any progeny thereof 35 containing the mutated LAT gene of the invention. More particularly, germ cells and somatic cells of said animals contain the mutated LAT gene of the invention as a result of

PCT/IB03/01044

chromosomal incorporation into the animal genome, or into an ancestor of said animal. Preferably, said mutated LAT gene is incorporated into the animal genome by targeted insertion (homologous recombination) in order to keep said mutated LAT 5 gene under the control of regulatory regions of the endogenous LAT gene. .

Therefore, a further object of the invention is to provide a mutated mouse gene coding for a mutant LAT protein, the 10 sequence of which corresponds to a wild type sequence and contains the single mutation of the tyrosine Y136, or a composite mutation of the tyrosine residues at positions 175, 195, and 235. Said mutation consists in the replacement by a residue preventing the association of the "tyrosinewith the SH2 domain of proteins. 15 based" sequences Preferably, said mutation of the tyrosine leads to its replacement by a phenylalanine, an aspartic acid or a glutamic acid. More preferably, said mutation of the tyrosine leads to its replacement by a phenylalanine. In a 20 preferred embodiment, the sequence of the gene encoding mutated mouse LATY136F protein corresponds to sequence ID N°1. The invention further includes chromosomal DNAs containing exon 7 of the mutated gene (SEQ ID N°2). The invention concerns a mouse containing said mutated mouse gene.

25

The present invention also encompasses plasmids comprising a DNA or a part thereof, having a sequence corresponding to the mutated gene according to the invention. In a preferred the plasmids of the invention contain embodiment, 30 restriction enzyme cleavage site, which is introduced in the intron 3' of exon 7. Advantageously, the restriction enzyme cleavage site is a Bgl II restriction site.

Said plasmids are useful for the generation of non-human 35 animals according to the present invention.

Consequently, the invention also encompasses non-human

embryos introduced with the plasmids of the invention, and non-human embryos obtained by homologous recombination using the plasmids of the invention. In a preferred embodiment, the non-human embryos are embryonic stem cells derived from a mouse. Advantageously, the ES cells are CK35 129/SV ES cells.

The invention also concerns the LAT mutant murine protein sequence containing the single mutation of the tyrosine Y136 10 or a composite mutation of the tyrosine residues at positions 175, 195, and 235. Said mutation consists in the replacement by a residue preventing the association of the "tyrosine-based" sequences with the SH2 domain of proteins. Preferably, said mutation of the tyrosine leads to its replacement by a phenylalanine, an aspartic acid or a glutamic acid. More preferably, said mutation of the tyrosine leads to its replacement by a phenylalanine. In one embodiment, the invention concerns a mutated LAT protein containing the mutated amino acid sequence of exon 7 (SEQ ID N°3).

The magnified and accelerated sequence of pathological events observed in the LAT Y136F, and LAT Y175F+Y195F+Y235F mice permits to readily start tests and studies. For example, 25 mutant LAT Y136F mice phenotype is achieved when they are 4 weeks old.

The mutant non-human animal according to the invention are useful in various applications of interest, in particular:

- 30 to analyze the impact of drugs on the molecular mechanisms that lead to exacerbated IgE production as well as tissue eosinophilia, and
- as a bioreactor allowing the dedicated production of IgE antibody of known specificity prior to or following a step
 35 of humanization of the mutated LAT mouse (preferably LAT*136F or LAT *175F+*195F+*235F mouse).

Consequently, the present invention provides models of allergy, and/or asthma disease comprising animals according to the invention. In particular, the animals of the invention can be used as models of eosinophilia and/or $T_{\rm H}2$ cells deregulation, more particularly $T_{\rm H}2$ cells accumulation.

PCT/IB03/01044

Therefore, the invention concerns the use of a mutant non-human animal according to the present invention as a model of allergy and/or asthma disease. The invention also concerns the use of a mutant non-human animal according to the present invention as a model of eosinophilia. More generally, the invention concerns the use of a mutant non-human animal according to the present invention as a model of $T_{\rm H}2$ cells deregulation, more particularly a model of $T_{\rm H}2$ to the secumulation.

Due to the increased sensitivity of population, health difficulties such as asthma or allergies are more frequent. The animals according to the invention are suitable models 20 to help the research in these domains.

Accordingly, the present invention provides a method of screening for a drug for treatment of allergy, asthma and/or disease associated with $T_{\rm H}2$ cell deregulation or activity comprising the step of subjecting the animals according to the invention, which are administered with the drug to a comparison with said animals, not administered with the drug.

- 30 More particularly, the invention concerns a method of screening of drugs for treatment of allergy, asthma and/or disease associated with $T_{\rm H}2$ cell deregulation or activity comprising the step of:
- administering a candidate drug to a non-human animal
 having a LAT gene coding for a mutant LAT protein according to the present invention;
 - 2) evaluating the effect of said drug on the symptom or sign

of allergy, asthma and/or disease associated with $T_{\rm H}2$ cell deregulation or activity; and

3) selecting the drug that reduces said symptom or sign.

In a preferred embodiment, said screening method uses non5 human animals not administered with drugs as control
experiments. In an other preferred embodiment, said effect
of said drug can be evaluated by measuring at least one
parameter selected from the group: IgE level, IgG1 level,
interleukin level (preferably IL-4, IL-10, IL-5 and/or IL10 13), and eosinophilia. More preferably, said effect of said
drug is evaluated by measuring the serum level of IgE and/or
IgG1.

The invention also contemplates a method of screening drugs 15 for treatment of allergy, asthma and/or disease associated with $T_{\rm H}2$ cell deregulation or activity comprising the step of:

- subjecting cells having a LAT gene coding for a mutant LAT protein according to the present invention to a 20 candidate drug;
 - 2) evaluating the effect of said drug on said cells;
 - 3) selecting the drug having the desired effect.

In a preferred embodiment, said effect of said drug can be evaluated by measuring the interleukin production, more 25 particularly the IL-4 production.

An other object of this invention resides in a method of screening drugs that regulate the activity of $T_{\rm H}2$ cells, comprising the step of:

- 30 1) administering a candidate drug to a non-human animal having a LAT gene coding for a mutant LAT protein according to the present invention; and
 - 2) selecting a drug that modulates the activity of $T_{\rm H}2$ cells in said non-human animal.

35

The screening methods can be used to select, identify, characterize and/or optimize candidate drugs. The candidate

WO 03/068968 PCT/I

PCT/IB03/01044

drugs may be of any origin, nature and structure. Their concentration may be adjusted by the skilled artisan. Furthermore, several drugs may be tested in parallel, or in combination.

5

A further object of this invention is a method of producing a pharmaceutical composition for treating a disease associated with deregulated $T_{\rm H}2$ cells activity, particularly asthma or allergy, the method comprising (i) selecting, identifying, optimizing or characterizing a compound using a screening assay as described above and (ii) conditioning said compound, or a derivative thereof, in a pharmaceutically acceptable carrier or vehicle.

15 In still another application, the present invention provides bioreactors for a large-scale production of human IgE antibodies comprising the animals according to the invention. LAT 136F and LAT 175F+195F+1235F mice are indeed able to produce tremendous amount of IgE, as it is described in 20 example 2. IgE produced by mutant mice are useful for applications such as desensitization or for kit of clinical assay.

Therefore, the invention concerns a method of production of 25 human IqE antibodies comprising the steps of :

- 1) providing a non-human animal expressing humanized IgE;
- 2) breeding said animal expressing humanized IgE with a non-human animal having a LAT gene coding for a mutant LAT protein according to the present invention;
- 30 3) immunizing the animal of the progeny with an allergen;
 - 4) recovering humanized IgE specific to said allergen.

The step 4 can comprise the step of producing B cell hybridomas producing said humanized IgE specific to said allergen. The invention relates to said B cell hybridoma

35 producing said humanized IgE specific to said allergen.

Said non-human animal expressing humanized IgE can be

obtained by conventional knock-in in which the genetic segment corresponding to the constant exons of the IgE gene is substituted by the corresponding human sequence.

- 5 The invention concerns the non-human animal resulting from the breeding of the animal expressing humanized IgE with the non-human animal having a LAT gene coding for a mutant LAT protein according to the present invention.
- 10 The produced humanized IgE specific to said allergen can be used for desensitization and in clinical assays aiming at characterizing allergens, preferably atopic allergens, present in patient.
- oligonucleotide probes invention contemplates the 15 The specific to a mutated LAT gene coding for a mutant LAT protein containing the single mutation of the tyrosine corresponding to Y136 in the mouse LAT protein or a composite mutation of the three distal tyrosines 20 (corresponding to Y175, Y195 and Y235 in the mouse LAT protein). Such probes are useful to detect the presence of the mutation in a LAT gene. Hence, the invention provide oligonucleotides, the sequence of which corresponds to SEQ ID N°4 and/or SEQ ID N°5 as probes to screen the presence of 25 the mutation Y136 in the mouse LAT gene. More particularly, the invention concerns oligonucleotide probes specific to a mutated human LAT gene coding for a mutant LAT protein containing a single mutation of the tyrosine Y132 or a composite mutation of the tyrosine residues Y171, Y191 and 30 Y226. Such probes are useful for the detection of mutant LAT gene involved in asthma, allergy, eosinophlia and/or any disease associated with a $T_{\rm H}2$ cells deregulation or activity. Said probes can be part of a diagnostic kit.
- 35 Therefore, the invention relates to a diagnostic method for asthma, allergy, eosinophilia and/or $T_{\rm H}2$ cells deregulation, more particularly $T_{\rm H}2$ cells accumulation, comprising the

13

WO 03/068968 PCT/IB03/01044

detection of a mutated LAT gene coding for a mutant LAT protein containing a single mutation of Y132 or a composite mutation Y171+Y191+Y226. Additionally, the invention concerns a diagnostic kit for asthma, allergy, eosinophilia and/or $T_{\rm H}2$ cells deregulation, more particularly $T_{\rm H}2$ cells accumulation, comprising oligonucleotide probes for the detection of a mutated LAT gene coding for a mutant LAT protein containing a single mutation of Y132 or a composite mutation Y171+Y191+Y226.

10

Other characteristics and advantages of the invention are given in the following examples with reference to figures 1 to 11.

15 EXAMPLES

Mutation LAT Y136

Example 1: Production of mutant mice

To test in vivo the importance of LAT^{Y136} , the inventors 20 generated knock-in mice with a mutation replacing Y136 with phenylalanine (Y136F).

1. Materials and methods

25 Mice

Mice were maintained in a specific pathogen-free animal facility.

LAT Y136F mutation.

30 LAT genomic clones were isolated from a 129/01a phage library. After establishing the nucleotide sequence and the exon-intron structure of the LAT gene, the tyrosine residue found at position 136 and encoded by exon 7 was mutated to phenylalanine. Mutagenesis was performed on a 1717-bp Eco 35 RI-Xba I fragment encompassing part of exon 5, exons 6, 7 and 8. In addition to the intended mutation, a new Bgl II

restriction enzyme cleavage site was introduced in the intron 3' of exon 7 to accommodate the LoxP-flanked neo gene and facilitate subsequent identification of LATY136F mutant mice. Finally, the targeting construct was extended to give 5 1.7 kb and 4.8 kb of homologous sequences 5' and 3' of the EcoRI-XbaI fragment, respectively (see Fig. 2). After electroporation of CK35 129/SV ES cells (C. Kress et al., 1998), and selection in G418, colonies were screened for homologous recombination by Southern blot analysis. The 5' 10 single-copy probe is a 0.9-kb Bgl II-Xba I fragment isolated from a LAT genomic clone. When tested on Bgl II-digested DNA, the 5' probe hybridizes either to a 8.5 kb wild-type fragment or to a 4.5 kb recombinant fragment. Homologous recombination events at the 3' side were screened by long 15 range PCR. Homologous recombinant ES clones were further checked for the presence of the intended mutation by sequencing the genomic segment corresponding to exon 7. Finally, a neo probe was used to ensure that adventitious non homologous recombination events had not occurred in the 20 selected clones.

Production of mutant mice.

Mutant ES cells were injected into Balb/c blastocysts. Two $\mathtt{LAT}^{\mathtt{Y136F}}$ recombinant ES cell clones were found capable of germ 25 line transmission. The two mutant mouse lines were first bred to Deleter mice (Schwenk. F et al., 1995) to eliminate the Lox P-flanked neomycin cassette, and intercrossed to produce homozygous mutant mice. The two independentlyderived mutant lines showed indistin-guishable phenotype. To 30 confirm that the LAT Y136F mutation had been genuinely introduced, LAT transcripts were cloned by transcription and PCR amplification from the thymus of the mutated mice, and the presence of the intended mutation confirmed by DNA sequence analysis. Screening of mice for 35 the presence of the LAT Y136F mutation was performed by PCR using the following pairs of oligonucleotides:

c : 5'-GTGGCAAGCTACGAGAACCAGGGT-3' (SEQ ID N°4);

d: 5'-GACGAAGGAGCAAAGGTGGAAGGA-3' (SEQ ID N°5).

The single Lox P site remaining in the LAT Y136F allele after deletion of the neo^r resulted in an amplified PCR product 140 bp-longer than the 510 bp-long fragment 5 amplified from the wild-type LAT allele.

2) Mutant mice development

Mice homozygous for the LAT^{Y136F} mutation, hereafter denoted LAT Y136F, were born at expected Mendelian frequencies and 10 their T cells contained levels of LAT proteins similar to wild-type T cells. At birth LATY136F mice displayed peripheral lymphoid organs of normal size. Beginning at about 3 weeks, however, the spleen and lymph nodes of the mutant mice started to enlarge relative to wild-type littermates, such weeks of age, spleen cellularity was 15 approximately 10 times that of wild-type mice (Fig. 3 A-C). Despite marked lymphocytic infiltrations in the lung, liver and kidney, homozygotes lived to at least 17 weeks of age, and no chronic intestinal inflammation or tumor formation 20 was observed. The effects of the LATY136F mutation were only detectable after breeding mice to homozygosity or to mice carrying a null allele of the LAT gene.

25 Example 2: Effect of the mutation: spontaneous exaggerated T helper type 2 immunity in mice

1. Materials and methods

30 Purification of CD4+ T cells and eosinophils.

Lymph node and spleen cells from several mice were pooled and the CD4+ cells purified using a high gradient magnetic cell separation system (S. Miltenyi et al., 1990). Eosinophils were sorted on a FACSvantageTM on the basis of their FSChigh, HSA+, and CD11b+ phenotype.

Antibodies and flow cytometric analysis.

Before staining, cells were preincubated on ice for at least 10 min with polyclonal mouse and rat Ig to block Fc receptors. Flow cytometric analysis was performed as 5 described previously (M. Malissen et al., 1995). All the antibodies were from BD PharMingen except the anti-CCR3 antibody that was purchased from R&D.

Staining for intracellular cytokines.

10 Before intracellular cytokine staining, cells (1.5×10^6) were cultured for 4 h in the presence of monensin (GolgiStop; BD PharMingen) at a final concentration of 2 µM. Cells were then immediately placed on ice, washed, resuspended in PBS 1X, 1% FCS, 0.20% sodium azide, and 15 stained with an APC-conjugated anti-CD4 antibody. intracellular cytokine staining, cells were first fixed using the cytofix/cytoperm kit (BD PharMingen). Each cell sample was subsequently split into aliquots that were separately stained with (1) a combination of FITC-conjugated 20 anti-IFN- α and PE-conjugated anti-IL-2 antibodies, (2) a combination of FITC-conjugated anti-IL-5 and PE-conjugated anti-IL-4 antibodies, and (3) a combination of fluorochromeconjugated and isotype-matched negative control Ig (BD PharMingen). After a final wash, CD4+ cells (104) were 25 analyzed on a FACSCalibur TM flow cytometer after gating out dead cells using forward and side scatters.

RNase protection assay.

For multiplex cytokine transcript analysis, total cellular 30 RNA was isolated from the specified cells using TRIzol (GIBCO-BRL Life Technologies) and analyzed by ribonuclease protection assay using an MCK-1 RiboQuantTM custom mouse template set (BD Pharmingen). Briefly, ³²P-labeled riboprobes were mixed with 10 μg of RNA, incubated at 56°C for 12 to 16 35 hours, and then treated with a mixture of RNases A and T1 and proteinase K. RNase-protected ³²P-labeled RNA fragments

were separated on denaturing polyacrylamide gels and the intensity of the bands evaluated with a Fuji imaging plate system.

5 Determination of serum isotype-specific immunoglobulin levels.

The titres of polyclonal IgM, IgG1, IgG2a, IgG2b, IgG3 and IgA antibodies and κ and λ light chains were determined using isotype-specific ELISA (Southern Biotechnology). The concentrations of IgG1 and IgE were determined by comparing test sample dilution series values with isotype control standards.

2. Results

15 A prominent phenotype of the CD4 T cells found in LAT 136F mice was revealed when the inventors measured their ability to make cytokines. Due to the short half-lives of cytokines and of their transcripts, their analysis generally requires restimulation of T cells in vitro with PMA and ionomycin. A 20 multiprobe RNase protection assay detecting levels of transcripts of 9 cytokines showed that CD4 T cells freshly isolated from LAT 136F mice contained sufficient IL-4 and IL-10 transcripts to allow their detection even without ex vivo restimulation (Fig. 4A). Upon activation by PMA/ionomycin 25 the levels of IL-4 and IL-10 transcripts they contained were further increased, and IL-5, IL-13, and IFN- α transcripts became readily detectable (Fig. 4B). In marked contrast, wild-type CD4 T cells yielded only the IL-2 and IFN- α transcripts expected for primary T cells. Analysis of IL-4 30 production at the single cell level, showed that following a 4 hr activation with PMA/ionomycin, close to 80% of the CD4 T cells isolated from LAT^{Y136F} mice expressed very high levels of IL-4 (Fig. 4C). Consistent with the notion that these CD4 T cells were refractory to TCR stimuli, none of them scored 35 as IL-4+ in response to TCR cross-linking (Fig. 4C). Thus, LAT Y136F spontaneously developed a high frequency of $T_{\rm H}2$

PCT/IB03/01044 WO 03/068968

cells. In the case of wild-type CD4 T cells, TH2 polarization of such magnitude is only achieved following prolonged antigenic stimulation in the presence of IL-4.

- 5 Light scatter analysis of thymic and lymph node cells from LATY136F mice older than 4 weeks revealed a unique cell population that was almost absent from age-matched wild-type mice, and showed both an intermediate forward scatter and a high side scatter (Fig. 5A, 5B, 6A). Based on several of 10 criteria, these cells were identified as eosinophils (Fig. 6). Minute numbers of eosinophils normally reside in wildtype thymi, and their augmentation in LAT 136F thymi may primarily result from an intrinsic expression of LATY136F molecules. However, LAT transcripts were undetectable in 15 eosinophils purified from LAT vi36F mice, meaning that the thymic and lymph node eosinophilia they manifest result from the production of IL-5 by the abnormal CD4 cells present in these mutant mice.
- 20 Secondary lymphoid organs of 6-week old LATY136F contained 7 to 10 times more B cells than their wild-type splenomegaly and generalized counterparts. Thus, the lymphadenopathy that developed in young LAT History mice can be mostly accounted for by cells belonging to the T and B cell 25 lineages. Over 90% of the mature B cells found in the spleen and lymph nodes of 6-week old wild type littermates had a resting phenotype (Fig. 7A). In marked contrast, only 25% of the B cells found in the enlarged secondary lymphoid organs LAT^{Y136F} littermates showed age-matched 30 phenotype. Among the remaining B cells, 25% showed an hyperactivated phenotype, and 50% expressed a phenotype typical of antibody producing cells. Coincident with the presence of these latter cells, serum IgG1 concentrations were elevated approximately 200 times compared to wild-type 35 mice, whereas those of IgE were elevated 2500 to 10000 times (Fig. 7C). In contrast, the levels of the other Ig isotypes did not differ significantly from those of wild-type serum

PCT/IB03/01044

WO 03/068968

(Fig. 7B). In support of a polyclonal hypergammaglobulinemia G1 and E, the concentrations of kappa and lambda light chains were both markedly augmented in the serum of LAT^{Y136F} IgG1 Notably, IgE and 7B). mice (Fig. 5 concentrations reached a plateau as early as 5 weeks of age (Fig. 7C), the values of which exceeded the extraordinarily large amounts of IgE and IgG1 previously reported for mice deprived of NFATc2 and NFATc3 transcription factors. Given that B cells do not express LAT proteins, and considering 10 that isotype switching to IgE and IgG1 is highly dependent on the presence of IL-4 and IL-13, the overproduction of IgE and IgG1 noted in LAT mice is secondary to the presence

15 Example 3: Production of IgE

Mice expressing humanized IgE are developed by conventional knock-in strategy in which the genetic segment corresponding to the constant exons of the IgE gene is substituted by the corresponding human sequence. Mice with a humanized IgE 20 locus are bred into LAT Y136F mice. Following immunization, B cell hybridomas producing specific human IgE are produced, and the resulting specific human IgE are used as "standard" in clinical assays aiming at characterizing atopic allergens present in patients."

of an abnormally high frequency of $T_{\rm H}2$ effectors.

25

Example 4: Screening for a drug

Mutant mice and control ones will be treated with a variety of drugs or original compounds. Their effects will be analyzed in vivo by measuring various parameters such as:

- TH2 cells differentiation. 30
 - Production of T_H2 types cytokines
 - Eosinophilia
 - Hypergammaglobulinemia G1 and E .



Mutation LAT*175+Y195+Y235

Example 5: Production of mutant mice

To test in vivo the importance of the three carboxy-terminal tyrosines (LAT Y175, LAT Y195 and LAT Y235), the inventors generated knock-in mice with a mutation replacing these three tyrosines with phenylalanine (LAT Y175F + Y195F + Y235F).

10 1. Materials and methods

Mice

Mice were maintained in a specific pathogen-free animal facility.

15

LAT Y175F + Y195F + Y235F mutation.

LAT genomic clones were isolated from a 129/01a phage library. After establishing the nucleotide sequence and the exon-intron structure of the LAT gene(EMBL Nucleotide 20 Sequence Datatase; accession number: AJ438435), the tyrosine residues found at positions 175, 195 and 235 and encoded by exons 9, 10, 11 were mutated to phenylalanine. Mutagenesis was performed on a 815 bp Ncol-BamHI fragment encompassing exons 9, 10, 11 (coding for tyrosines 175 (exon 9), 195 and 235 (exon 11) and part of 25 (exon 10) exon . 12 (corresponding to the 3' untranslated region of LAT). Each exon was mutated independently and new restriction sites were introduced for facilitating subsequent cloning steps. A new Eco RI site was introduced on the 5' side of the NcoI 30 site, BamHI and ClaI sites were introduced between exons 9 and 10, a HindIII site was introduced between exons 10 and 11, and BqlII, XhoI, and NotI sites were introduced on the 3' side of exon 11 in lieu of the original BamHI site. PCR reactions were performed with Pwo DNA polymerase (Boehringer 35 Mannheim), and PCR products were purified and cut with with EcoRI and BamHI for exon 9, BamHI and HindIII for exon 10,

and HindIII and NotI for exon 11. These three fragments were assembled in a pBS-KS plasmid (Stratagene). The resulting plasmid was used to clone a 3.5 kb Eco RI-Nco I genomic fragment providing a 5' homology arm and a 4.3 kb-Sal I 5 genomic fragment providing a 3' homology arm. Finally a loxP flanked neor gene was introduced using the BamHI and ClaI sites that were engineered between exons 9 and 10. After electroporation of CK35 129/SV embryonic stem (ES) cells and selection in G418, colonies were screened for homologous 10 recombination by southern blot analysis using a 3' singlecopy probe that consisted of a 1.1 kb EcoRI-HindIII fragment isolated from a LAT genomic clone. When tested on BamHI digested genomic DNA, the 3' probe hybridizes either to a 7.0 kb wild-type fragment or to a 9.1 kb recombinant 15 fragment. The presence of a genuine recombination event was checked by PCR using the following pair of primers (depicted in Figure 8):

- f: 5'-CCCAGAGGCAAACCCTCTGAAG-3' (SEQ ID N°6) and
- g: 5'-TCGAATTCGCCAATGACAAGACGC-3'(SEQ ID N°7). This PCR 20 gives a band of 8.6 kb in the recombinant ES clones only. Homologous recombinant ES clones were further checked for the presence of the intended mutations by sequencing the genomic segment corresponding to exons 9, 10 and 11. Finally, a neo probe was used to ensure that adventitious 25 non-homologous recombination events had not occured in the selected clones.

Production of mutant mice.

Mutant ES cells were injected into Balb/c blastocysts. Two
30 LAT Y175F + Y195F + Y235F recombinant ES cell clones were
found capable of germ line transmission. The two mutant
mouse lines were first bred to Deleter mice (Schwenk. F et
al., 1995) to eliminate the Lox P-flanked neomycin cassette,
and intercrossed to produce homozygous mutant mice. The two
35 independently-derived mutant lines showed indistinguishable
phenotype. To confirm that the LAT Y175F + Y195F + Y235F
mutation had been genuinely introduced, LAT transcripts were





cloned by reverse transcription and PCR amplification from the thymus of the mutated mice, and the presence of the intended mutation confirmed by DNA sequence analysis. Screening of mice for the presence of the LAT Y175F + Y195F 5 + Y235F mutation was performed by PCR using the following pairs of oligonucleotides:

d:5'-GGAGACTTAGATGTCTGAGCCG-3' (SEQ ID N°8) and

- e: 5'-GACAGACCAGCAGGGACAGTG-3' (SEQ ID N°9) (Wt 238bp, mutant 435bp).
- 10 The single Lox P site remaining in the LAT Y175F + Y195F + Y235F allele after deletion of the neo^r resulted in an amplified PCR product 140 bp-longer than the 510 bp-long fragment amplified from the wild-type LAT allele.

15 2) Mutant mice development

Mice homozygous for the LAT Y175F + Y195F + Y235F mutation, hereafter denoted LAT Y175F + Y195F + Y235F were born at expected Mendelian frequencies and their T cells contained levels of LAT proteins similar to wild-type T cells. At 20 birth LAT Y175F + Y195F + Y235F mice displayed peripheral lymphoid organs of normal size. Beginning at about 3 months, however, the spleen and lymph nodes of the mutant mice started to enlarge relative to wild-type littermates, such by 3 months of age, spleen cellularity was that 25 approximately 5 times that of wild-type mice. Homozygotes lived to at least 5 months of age, and no chronic intestinal inflammation or tumor formation was observed. The effects of the LAT Y175F + Y195F + Y235F mutation were only detectable after breeding mice to homozygosity or to mice carrying a 30 null allele of the LAT gene.

Example 6: Effect of the mutation: a subset of gamma/delta T cells expands and acquire a spontaneous exaggerated T helper type 2 immunity in mice.



Purification of gamma/delta T cells and eosinophils.

Spleen cells from several mice were pooled and the gamma/delta T cells purified using a high gradient magnetic 5 cell separation system (S. Miltenyi et al., 1990).

Antibodies and flow cytometric analysis.

Before staining, cells were preincubated on ice for at least 10 min with polyclonal mouse and rat Ig to block Fc receptors. Flow cytometric analysis was performed as described previously (M. Malissen et al., 1995). All the antibodies were from BD PharMingen.

Staining for intracellular cytokines.

Before intracellular cytokine staining, cells (1.5×10^6) 15 were cultured for 4 h in the presence of monensin (GolgiStop; BD PharMingen) at a final concentration of 2 µM. Cells were then immediately placed on ice, resuspended in PBS 1X, 1% FCS, 0.20% sodium azide, and stained with an APC-conjugated anti-CD5 antibody. 20 intracellular cytokine staining, cells were first fixed using the cytofix/cytoperm kit (BD PharMingen). Each cell sample was subsequently split into aliquots that were separately stained with (1) a combination of FITC-conjugated anti-IFN-D and PE-conjugated anti-IL-2 antibodies, (2) a 25 combination of FITC-conjugated anti-IL-5 and PE-conjugated anti-IL-4 antibodies, and (3) a combination of fluorochromeconjugated and isotype-matched negative control Ig PharMingen). After a final wash, CD5+ cells (104) were analyzed on a FACSCalibur™ flow cytometer after gating out 30 dead cells using forward and side scatters.

RNase protection assay.

For multiplex cytokine transcript analysis, total cellular RNA was isolated from the specified cells using TRIzol 35 (GIBCO-BRL Life Technologies) and analyzed by ribonuclease protection assay using an MCK-1 RiboQuantTM custom mouse



template set (BD Pharmingen). Briefly, ³²P-labeled riboprobes were mixed with 10 µg of RNA, incubated at 56°C for 12 to 16 hours, and then treated with a mixture of RNases A and T1 and proteinase K. RNase-protected ³²P-labeled RNA fragments were separated on denaturing polyacrylamide gels and the intensity of the bands evaluated with a Fuji imaging plate system.

Determination of serum isotype-specific immunoglobulin 10 levels.

The titres of polyclonal IgM, IgG1, IgG2a, IgG2b, IgG3 and IgA antibodies and □ and □ light chains were determined using isotype-specific ELISA (Southern Biotechnology). The concentrations of IgG1 and IgE were determined by comparing 15 test sample dilution series values with isotype control standards.

2. Results

A prominent phenotype of the CD90.2+, CD5+ gamma/delta T 20 cells found in LAT Y175F + Y195F + Y235F mice (see Figure 9) was revealed when the inventors measured their ability to make cytokines. Due to the short half-lives of cytokines and of their transcripts, their analysis generally requires restimulation of T cells in vitro with PMA and ionomycin. A 25 multiprobe RNase protection assay detecting levels transcripts of 9 cytokines showed that gamma/delta T cells freshly isolated from LAT Y175F + Y195F + Y235F mice contained large amounts of IL-4, IL-5, IL-10 and IL-13 transcripts to (Figure 10). This attribute is reminiscent of 30 the observation made with the alpha/beta T cels present in the periphery of the LAT Y136F mice. In marked contrast, wild-type CD4 T cells yielded only the IL-2 and IFN-γ transcripts expected for primary T cells. Analysis of IL-4 production at the single cell level, showed that following a 35 4 hr activation with PMA/ionomycin, close to 80% of the CD4 T cells isolated from LAT Y175F + Y195F + Y235F mice expressed very high levels of IL-4. Thus, LAT Y175F + Y195F

PCT/IB03/01044

+ Y235F mice spontaneously developed a high frequency of gamma/delta T cells with a Th2 phenotype. In the case of wild-type CD4 T cells, $T_{\rm H}2$ polarization of such magnitude is only achieved following prolonged antigenic stimulation in 5 the presence of IL-4.

The spleen of 3-month old LAT Y175F + Y195F + Y235F mice contained 5 to 10 times more B cells than their wild-type counterparts. Thus, the splenomegaly that developed in LAT 10 Y175F + Y195F + Y235F mice can be mostly accounted for by cells belonging to the T and B cell lineages. Over 90% of the mature B cells found in the spleen and lymph nodes of 3month old wild type littermates had a resting phenotype (Fig. 11A). In marked contrast, only 16% of the B cells 15 found in the enlarged secondary lymphoid organs of agematched LAT Y175F + Y195F + Y235F littermates showed a resting phenotype. Among the remaining B cells, 21% showed an hyperactivated phenotype, and 63% expressed a phenotype typical of antibody producing cells. Coincident with the 20 presence of these latter cells, serum IgG1 concentrations were elevated approximately 100 times compared to wild-type mice, whereas those of IgE were elevated 500 to 5000 times (Fig. 11). In contrast, the levels of the other Ig isotypes did not differ significantly from those of wild-type serum. 25 Given that mature B cells do not express LAT proteins, and considering that isotype switching to IgE and IgG1 is highly dependent the presence of IL-4and IL-13, on overproduction of IgE and IgG1 noted in LAT Y175F + Y195F + Y235F mice is secondary to the presence of an abnormally 30 high frequency of gamma/delta T cells producing cytokines.

Example 7: Production of IgE

Mice expressing humanized IgE are developed by conventional 35 knock-in strategy in which the genetic segment corresponding to the constant exons of the IgE gene is substituted by the corresponding human sequence. Mice with a humanized IgE



locus are bred into LAT Y175F + Y195F + Y235F mice. Following immunization, B cell hybridomas producing specific human IgE are produced, and the resulting specific human IgE are used as "standard" in clinical assays aiming at 5 characterizing atopic allergens present in patients."

Example 8: Screening for a drug

Mutant mice and control ones will be treated with a variety of drugs or original compounds. Their effects will be 10 analyzed in vivo by measuring various parameters such as:

- $T_{\rm H}2$ cells differentiation.
- \bullet Production of $T_{H}2$ types cytokines
- Hypergammaglobulinemia G1 and E .



References

Kress, C., Vandormael-Pournin, S., Baldacci, P., Cohen-Tannoudji, M., and Babinet, C. (1998). Nonpermissiveness for mouse embryonic stem (ES) cell derivation circumvented by a single backcross to 129/Sv strain: establishment of ES cell lines bearing the Omd conditional lethal mutation, Mamm Genome 9, 998-1001.

Lin, J., and Weiss, A. (2001). Identification of the minimal 10 tyrosine residues required for linker for activation of T cell function, J Biol Chem 276, 29588-29595.

Malissen, M., Gillet, A., Ardouin, L., Bouvier, G., Trucy, J., Ferrier, P., Vivier, E., and Malissen, B. (1995).

15 Altered T cell development in mice with a targeted mutation of the CD3- epsilon gene, Embo J 14, 4641-53.

Miltenyi, S., Muller, W., Weichel, W., and Radbruch, A. (1990). High gradient magnetic cell separation with MACS, 20 Cytometry 11, 231-8.

Samelson, L.E., Bunnell, S.C., Trible, R.P., Yamazaki, T., and Zhang, W. (1999). Studies on the adapetr molecule LAT, Cold Spring Harbor Symposia On Quantitative Biology, Biology 25 Laboratory, Cold Spring Harbor, NY, n°64, 259-263.

Schwenk, F., Baron, U., and Rajewsky, K. (1995). A cretransgenic mouse strain for the ubiquitous deletion of loxP-flanked gene segments including deletion in germ cells, 30 Nucleic Acids Res 23, 5080-1.

Sommers, C.L., Menon, R.K., Grinberg, A., Zhang, W., Samelson, L.E., and Love, P.E. (2001). Knock-in mutation of the distal four tyrosines of linker for activation of T cells blocks murine T cell development, J Exp Med, 2001, 135-142.

28

WO 03/068968



Zhang, W., Trible, R.P., Zhu, M., Liu, S.K., McGlade, C.J., and Samelson, L.E. (2000). Association of Grb2, Gads, and phospholipas $C-\gamma 1$ with phosphorylated LAT tyrosine residues, J Biol Chem, 275, 23355-23361.



Claims

1. A non-human animal having a mutated LAT gene coding for a mutant LAT protein, wherein said mutant LAT protein leads to an exaggerated $T_{\rm H}2$ cell differentiation.

5

2. The non-human animal according to claim 1, wherein the sequence of said mutant LAT protein corresponds to a wild type sequence and contains a single mutation of the tyrosine corresponding to Y136 in the mouse LAT protein.

- 3. The non-human animal according to claim 2, wherein said mutated LAT gene coding for a mutant LAT protein comprises exon 7 of the mutated gene (SEQ ID No 2).
- 15 4. The non-human animal according to claim 1, wherein the sequence of said mutant LAT protein contains a composite mutation of the three distal tyrosine residues.
- 5. The non-human animal according to any of claims 1 to 4, 20 wherein said non-human animal is a mammal.
 - 6. The non-human animal according to claim 5, wherein said mammal is a rodent.
- 25 7. The non-human animal according to claim 6, wherein said rodent is a mouse.
- 8. The non-human animal according to any of claims 1 to 7, wherein said mutation consists in the replacement of the 30 tyrosine by a residue preventing the association of the "tyrosine-based" sequences with the SH2 domain of proteins.
- 9. The non-human animal according to any of claims 1 to 7, wherein said single mutation consists in the replacement of the tyrosine by a phenylalanine (Y-F), an aspartic acid (Y-



- D) or a glutamic acid (Y-E).
- 10. The non-human animal according to claim 9, wherein said single mutation consists in the replacement of the tyrosine 5 by a phenylalanine (Y-F).
 - 11. The non-human animal according to any of claims 1 to 10, wherein said non-human animal is homozygous for the mutated LAT gene or carries a null allele of the LAT gene.

10

- 12. The non-human animal according to any of claims 1 to 11, wherein said mutated LAT gene is incorporated into the animal genome by targeted insertion in order to keep said mutated LAT gene under the control of regulatory regions of the endogeneous LAT gene.
 - 13. A germ cell or somatic cell from a non-human animal according to any one of claims 1-12 or any progeny thereof containing the mutated LAT gene.

- 14. Use of an animal according to any of claims 1 to 12 as a model of allergy, asthma, eosinophilia and/or disease associated with $T_{\rm H}2$ cell deregulation.
- 25 15. A method of screening for a drug for treatment of allergy, asthma and/or disease associated with $T_{\rm H}2$ cell deregulation comprising the step of subjecting animals according to any of claims 1 to 12 which are administered with the drug to a comparison with said animals, not 30 administered with the drug.
 - 16. A method of screening for drugs for treatment of allergy, asthma and/or disease associated with $T_{\rm H}2$ cell deregulation comprising the step of:
- 35 1) administering a candidate drug to a non-human animal according to any of claims 1 to 12;
 - 2) evaluating the effect of said drug on the symptom or sign





of allergy, asthma and/or disease associated with $T_{\rm H}2$ cell deregulation; and

- 3) selecting the drug that reduces said symptom or sign.
- 5 17. The method according to claim 16, wherein said effect of said drug can be evaluated by measuring at least one parameter selected from the group of IgE level, IgG1 level, interleukin level (IL-4, IL-10, IL-5 and/or IL-13), and eosinophilia.

- 18. The method according to claim 17, wherein said effect of said drug can be evaluated by measuring the serum level of IgE and/or IgG1.
- 15 19. A method of screening drugs for treatment of allergy, asthma and/or disease associated with $T_{\rm H}2$ cell deregulation comprising the step of:
 - 1) subjecting cells according to claim 13 to a candidate drug;
- 20 2) evaluating the effect of said drug on said cells;
 - 3) selecting the drug having the desired effect.
 - 20. A method of screening for drugs for that regulate the activity of $T_{\rm H}2$ cells, comprising the step of:
- 25 1) administering a candidate drug to a non-human animal according to any of claims 1 to 12; and
 - 2) selecting a drug that modulates the activity of $T_{\rm H}2$ cells in said non-human animal.
- 30 21. A method of producing a pharmaceutical composition for treating a disease associated with deregulated $T_{\rm H}2$ cells activity, particularly asthma or allergy, the method comprising (i) selecting, identifying, optimizing or characterizing a compound using a screening method according
- 35 to any of claims 16 to 20 and (ii) conditioning said compound, or a derivative thereof, in a pharmaceutically acceptable carrier or vehicle.



22. A bioreactor for a large scale production of human IgE antibodies comprising an animal according to any of claims 1 to 12.

5

- 23. A method of production of human IgE antibodies comprising the steps of :
- 1) providing a non-human animal expressing humanized IgE;
- 2) breeding said animal expressing humanized IgE with a non-
- 10 human animal according to any of claims 1 to 12;
 - 3) immunizing the animal of the progeny with an allergen;
 - 4) recovering humanized IgE specific to said allergen.
- 24. The method according to claim 23, wherein the step 4 15 comprises the step of producing B cell hybridomas producing said humanized IgE specific to said allergen.
 - 25. A B cell hybridoma obtained by the method according to claim 24.

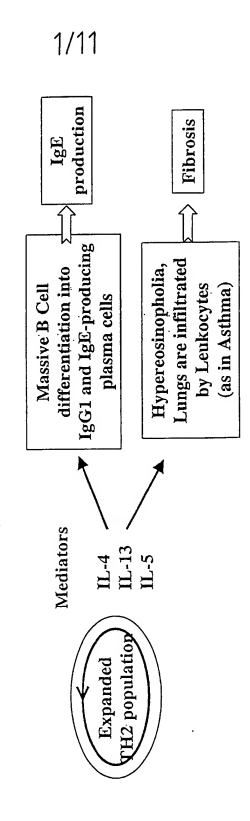
- 26. A mutated mouse gene coding for a mutant LAT protein, the sequence of which corresponds to a wild type sequence and contains a single mutation of the tyrosine Y136.
- 25 27. A mutated mouse gene coding for a mutant LAT protein, the sequence of which corresponds to a wild type sequence and contains a composite mutation of the three distal tyrosine residues.
- 30 28. The mouse gene according to any of claims 26 to 27, wherein said mutation consists in the replacement of the tyrosine by a phenylalanine (Y-F), an aspartic acid (Y-D) or a glutamic acid (Y-E).
- 35 29. The mouse gene according to 28, wherein said mutation consists in the replacement of the tyrosine by a phenylalanine (Y-F).



- 30. The mouse gene according to 26, wherein the sequence corresponds to sequence ID $N^{\circ}1$.
- 5 31. The mouse gene according to 30, wherein the sequence contains exon 7 of the mutated gene (SEQ ID $N^{\circ}2$).
- 32. A diagnostic method for asthma, allergy, eosinophilia and/or $T_{\rm H}2$ cells deregulation comprising the detection of a 10 mutated LAT gene coding for a mutant LAT protein containing a mutant LAT protein containing a single mutation of the tyrosine Y132 or a composite mutation of the three distal tyrosines Y171, Y191 and Y226.
- 15 33. A diagnostic kit for asthma, allergy, eosinophilia and/or $T_{\rm H}2$ cells deregulation comprising oligonucleotide probes for the detection of a mutated LAT gene coding for a mutant LAT protein containing a single mutation of the tyrosine Y132 or a composite mutation of the three distal 20 tyrosines Y171, Y191 and Y226.
 - 34. A non-human animal resulting from the breeding of a non-human animal expressing humanized IgE with the non-human animal according to any of claims 1 to 12.



igure 1





2/11

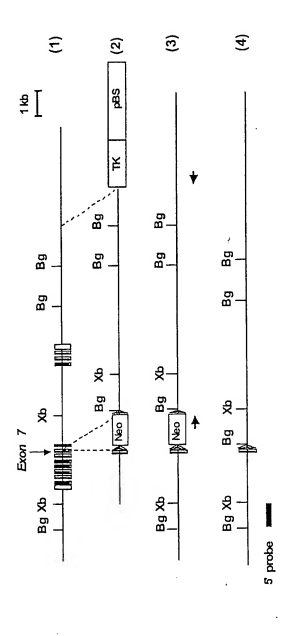


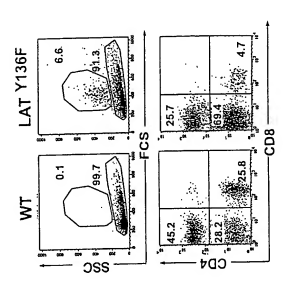
Figure (

Т**А**Ј **3**817 ΤW TAJ 4961Y ΤW M ΤW **436FY** TAJFigure 3

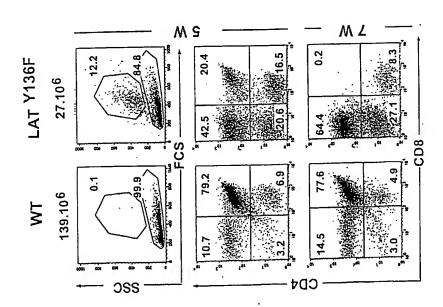


4/11 None LAT Y136F 9-71 ¥ \mathbf{m}

5/11



22



igur



CD11b

CD11c

CD11b

CD11c

CD

2

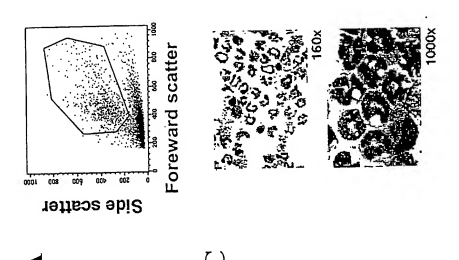
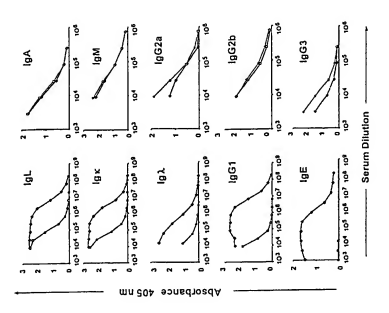
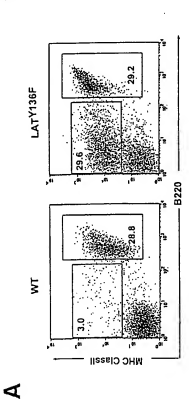


Figure 6

7/11



 $\mathbf{\omega}$



> 103 103

(Im\eq) f2el \$ \$ \$ \$

()

igur



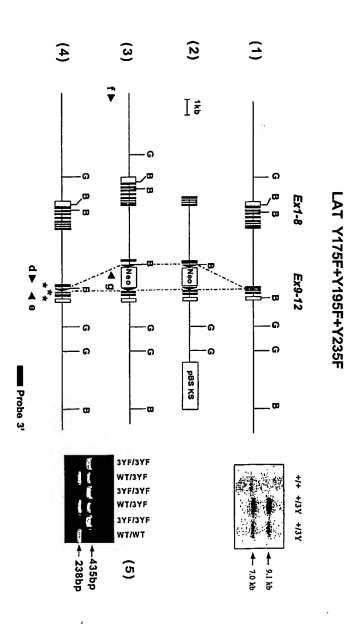


FIGURE 8



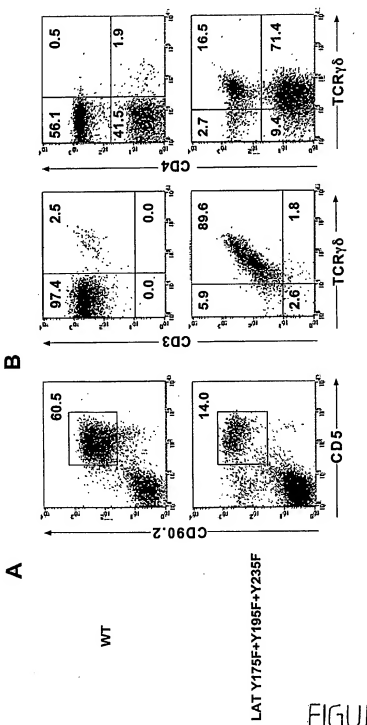


FIGURE 9





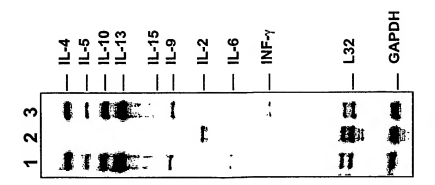


FIGURE 10



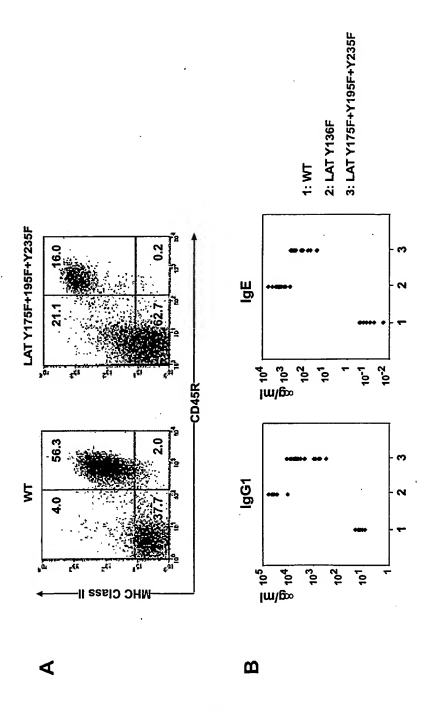


FIGURE 11





1 SEQUENCE LISTING

<110> INSTITUT NATIONAL DE LA SANTE ET DE LA RECHERCHE MEDICALE

<120> Mutated gene coding for a LAT protein and the biological applications thereof.

<130> B0184PCT

<140>

<141>

<150> US60/356.136

<151> 2002-02-14

<150> EP02/290610.1

<151> 2002-03-11

<160> 5

<170> PatentIn version 3.1

<210> 1

<211> 6307

<212> DNA

<213> Mus musculus

<400> 1
tatccatagt cccagactta acaggggctg tcaggtcacc ctgtgggtaa gtccctgtct 60
tctgagcttg gtaatctaga aggagggctg ctctttctg agtgagctgg ttcagtatga 120
ctgtgactca ccgtggtccc ctggaagtcg ctctcccagt agttaagcct gggagctggg 180
ggcctgtggt gccctcagtg ccctcggtcc acacaggect tggcagagcc tccttccagt 240
tctcccaccc gggcatgggg agggtaccgc gggcctggtt ggcacgtgtc tccttccta 300
gtggacgggc tgcctcatcc tgcagcctta gacccttcct ccacacagtc cctctgcctc 360

WO 03/068968



ctccccttc ccacaactgg gtgggggtga gtgggcaggg ggcaggctca gcctgctgag 420 cagectgatg atttectgee etcaceaeag etteetgteg caegeggtgg tgageaggag 480 aggcaggcgg ggagcaagaa aggggcaggt acagctgggc acggggatcg tgcagctggt 540 600 agctggggca cgggccccag ctctggctct ggggcgagca cctttccaga gccaacactg 660 ctctcaactc agtccagcaa gagagggag ccatccagcc ccgaaaggat acggctgcct 720 actgeeggge ggateceagg etggageeeg ettggteeea taeceetget gecactetgt 780 ctcqaqqqqc tqcaqtqcaq caqqqcctqt qgcaqqtqct ctgcaqatgg aagcagacgc cttgagcccg gtggggctgg gcctcctgct gctgcccttc ttggtcacgc tcctggctgc 840 900 cctqtqcqtq cqctqccqtq agttqccaqq taagtqggaa gctttqcgga actggatgat gggtgggcgc tecattggat ceteatacec tececagece etgeactete caetgteeet 960 1020 acctgggccc tgattgatgg tgggggcct gagtttcttt gtccctggtg caccccgatc 1080 ctgacttgtt ggatttcttt cctccagtct cctatgacag cacttccaca gagaggtgag tgggaagccc gtgtccctgt gtgtcttccc ttggttccac tcaagggttt ggggctgggg 1140 1200 ccctcttqqc cctqtaccca aqctqtctct ttcctqccaq tttqtaccca agaagcatcc 1260 tcatcaagcc acctcgtgag ttcagtgtct ctggccctcc tcgagggttt ttaagagtgt qcqtttqtcc ttqttcacct ttagctqtct gaagggctqt tccctqgctt gggatgggga 1320 aagtgggagc ccccatgtct gtctagggca tgttattttg gggtccattt gtccttcgag 1380 gccttgatgg ggggtgtctg gagccatccc tcaagcttca ttctgtgtcc tcagaaataa 1440 1500 ecgteeceg aacacetget gttteetace etetagteac tteetteeca eccetgagge agccagacet getececate cegtgagtat eccecaatte egtecettgg gtetactgtg 1560 cctctccacc ttctaggttg gggaggcgct ttttcctggt tgtcttgctc ccagagtcct 1620 1680 acctagacgt aatctctgac ctttggcttc caggagatcc ccacagcccc ttgggggttc 1740 ccatcggatg ccatcttccc agcagaattc agatgatggt aagggtgtag ggcacaggag ggctttgggg aggatgtaca acctgagctg atccagtctt cttctccctc tctctttgaa 1800 gccaacagtg tggcaagcta cgagaaccag ggtgggtctg gggtctgggg tagtgggtgg 1860 1920 qqtqqqqagg ctqqacctqt ccaqqtcqtq ttaactctcc tttctcacag agccagcctq taagaatgtg gatgcagatg aggatgaaga cgactatccc aacggcttcc tgtgagtggg 1980 tagaggagat etgacegtgg aagttgtgtg ceetttatea aettetegtt cetteettte 2040 2100 ttccagagtg gtgctgcctg acagtagtcc tgctgccgtc cctgttgtct cctctgctcc tqtqcctagc aaccctgacc ttgqaqacaq tqccttctct ggtqaqtcag gctttctgtc 2160 tacctccctc tgccatgtgc tgccagctct ccactcttgc ctccctctca cctccgtgac 2220

WO 03/068968 PCT/IB03/01044

			3			
gattgccgcc	cttccatttc	ctcctgtaga	cgttgggctt	cctgctcctc	atcacttccg	2280
actgtcttgt	ttttccttcc	acctttgctc	cttcgtctct	gttgtctaag	aaatttcctg	2340
actcttttg	aaccctgcca	ttgaaatttc	atttctcggc	tgggtgtgag	ggcctacgat	2400
cccagcatca	ggaggcagtg	gcaggagggt	tgaatttgag	gctagcctgg	gctacatagt	2460
gataccctct	cttcgaaaac	caaaacagca	cgacgatcaa	caaaaagaaa	acaaaagaat	2520
ttatttctct	tatctgaaag	teccettece	cttttttggc	gtctcggttc	tttttgtata	2580
gtacactgtt	gtttcttgga	agcaatatca	tctaatgtat	ctataagaac	tttgattaca	2640
tagccgggtg	gtggtggcgc	acgcctttaa	ttccagcact	cgggaggcag	aggcaggcgg	2700
atttctgagt	tcaaggccag	cctggtctac	agagtgagtt	ccaggacagc	caggactaca	2760
cagagaaacc	ctgtctcgaa	aaaacaaaac	aaaacaaatt	ttgattacag	attgtttctc	2820
tctgtgtctc	tatccctctc	tggttctgcc	cgtctctctg	tatctctgcc	cgtctctctg	2880
tatctctgcc	cgtctctctg	tatctctgcc	cgtctctctg	tatctctgcc	cgtctctctg	2940
tatctctgcc	cgtctctctg	tatctatctc	tgcccgtctc	tctgtatctc	tgcccgtctc	3000
tctgtatctc	tgcccgtctc	tctgtatctc	tgcctgtctc	tctcacacac	actcactgaa	3060
gatttattct	gcgtaccaca	tggtcgttgt	ttctcttggg	ctgcttttct	ctgctttggt	3120
ctttctcctt	ccttgagctt	ttctcaagtt	ctggtgatct	tcagttttct	atcctcttat	3180
ctctgtatag	catgagtatc	ccttacctga	aacacttcaa	tacagatttg	ggaatattta	3240
taaacatata	ataaattctc	ttggggatga	aactcaagat	aaaacatgta	attaatttat	3300
tcatgtttta	tacaaaccat	atatgtaata	tatacacagt	ctgaagatag	gtttttgttt	3360
tgtcttagtt	ttattggcat	agagcgtcat	tgtatagtcc	tggctgtcct	ggaacttgat	3420
atctagacca	ggtagactca	aactcaaatt	aaacgtgtag	gttaccatgc	tcggtcttta	3480
aggtagttct	atgcaaattt	taattaatct	tttgtatgaa	atagaagttt	catgaaattt	3540
tccatttgtg	gtatcgcacc	agtatgaaaa	ggttttggat	ttcggaatat	gatgaatttt	3600
ggagttttaa	aaggaacacc	caaccttctg	tatttaccct	agactattat	gtctgtactc	3660
tggctctgtt	ttgtttgaga	gagaatctca	ctgtagagtc	ctggctgccc	tggaactcac	3720
tttgtagatt	aagtatggcc	tttaactcca	gttgcctctg	gcttctgagt	tctgggatta	3780
tatggggtta	aagacgtatc	cctcttgttc	cacttggttt	ttgttgttgg	tggtttgttt	3840
atttagcttt	tttttttca	gtttttctcc	ctcaatacag	cttttctcta	tgtatccttg	3900
gctgtcctag	acctcactct	gtagaccagg	ctgtccttga	actcagaaat	ctgcctgcct	3960
ctacctcctg	agtgctggga	ttaaaggcac	gtgccaccac	cacctggctc	tcttgctcca	4020
tttgtaaccc	actgactata	caatgagtcc	ccatgtcaat	aaaaccaaga	caaaacaaaa	4080
acctagcttc	agactgcgta	tatatgattt	atataaacca	tgcatgactt	aattccgtgt	4140

WO 03/068968 PCT/IB03/01044

aatttgtcat	ttctctcctg	aaccccagac	tgtttgagtg	atcccttcct	tecateegte	4200
ctggtctctc	gctcctcatt	tcctggttat	gtctgctgac	ttttgctagg	gatttaggga	4260
gccaatgcag	caaacttgta	atggtaaaag	gatcattgct	aggggcaaaa	tgactcattt	4320
taatttcagt	gagagactct	gtctcaaaga	actatggtgg	aatggctaaa	gcctccatgt	4380
gctcctgagt	gtgtgcagtg	gcataacaca	cagagaggta	ctaagagaac	tactgttaac	4440
tgaggagcaa	ctctatgccc	tcgtggtgtg	tacagctcat	tagacctcac	agttcgtggg	4500
tgctctgctg	accgtaccct	cttcccctcc	tgtccctcac	atctctctct	gtactgtctc	4560
tctgtatggt	atgctagagt	ttatttattt	acttaaattg	atacagtctt	gctgttgtga	4620
tgtccagtct	gtcttgagct	caagttagcg	cctgcctccc	gtcttctgag	accacagcct	4680
ggctcaaggt	tgctagtaat	tggaacaacg	gtagcacata	gtgtattgca	ggctctgttt	4740
tacaatttat	tgtttattcc	tcactctagt	ccttccaggc	aggtcctgtt	atgaacctca	4800
ttctacagac	taggaaactg	gggcagggag	catttaggtg	acttatctga	ggttagatag	4860
ttgcttagtg	ctgggactga	ggtttgagcc	agtgtatttg	gctcagcttg	tccacatgcc	4920
catacagaaa	ccaggcaacc	atgaaaccag	aaagcaaaaa	gctgtgtagc	attgtgagtg	4980
acctttgtgg	gcccaggaag	gtgagggcaa	gagctgataa	cattgagaga	ccaacaggtc	5040
tgagaagagg	ggatgccaac	tagaccaagt	gtgccacttc	ttcacagatc	accaaggtct	5100
ctgcactctg	agctccttgg	agccctgctc	tccagcctca	ctgcctgagt	cctgtattgt	5160
ctctgttcca	ttcccccaga	ggctctggtc	ctggctctcc	atccacctcc	atggcccttg	5220
ccctgcccag	gcttcttctc	ccctcgcttt	tcctgaatat	tctctctata	ttgtgagtct	5280
gcctgggggt	tgtgttagga	gacttagatg	tctgagccgg	gggtgggagg	tgtctctggg	5340
gaacagtgcc	tggctgagtg	tctgctaata	actgtactgc	aatggctatt	ctacagtgga	5400
gtcgtgtgaa	gattacgtga	atgttcctga	gagtgaggag	agcgcagagg	cgtctctggg	5460
taggtgactc	tgcactccat	gcatggccca	tageetetee	ctaccctctg	catggcctgc	5520
ccttcacacc	actgtccctg	ctggtctgtc	cccacagatg	ggagccggga	gtatgtgaat	5580
gtgtccccag	agcagcagcc	agtgaccagg	gctgagctgg	gtgagtacca	aggtgtaagg	5640
gggcagaggc	tggggagcag	ccttgagtag	agagtctgta	ggctgaacgg	cagteteect	5700
ctgtttttcc	ctctcagcct	ctgtgaactc	ccaggaggtg	gaagacgaag	gagaagagga	5760
	ggagaggaag					5820
tagtgagtgg	tctctgtccc	cgcccccacc	ttgggccttc	tetecaggae	cccctcctg	5880
	gtggttaggc		•			5940
tgtcctccct	gtctcccctg	ccctgctgtg	tttcagctgc	agctgtctgt	cctgaaactg	6000



WO 03/068968

5 6060 gacttgctgg ggtgtcgcta agaggatccc atttgacctc tgccttgcca cagcctgaga 6120 atcttcccct aacttattgt cactttgggg tccagtctgt gtccccaata ttctgtacct tctgataaag cctgagaatg aatctggttc cagccagacc atgtcatgga ataaaggcca 6180 6240 tgtgacataa agtcgtcgtt gtcttctttt tgttgttgct ggtgttgttg gtttgttgt 6300 ttgtttaact gggacagggt cttgctatgt tgatcaaggc tggtcttgaa cctgtgggtg 6307 atcatcc <210> 2 <211> 61 <212> <213> Mus musculus <400> 2 agccagcctg taagaatgtg gatgcagatg aggatgaaga cgactatccc aacggcttcc 60 61 t <210> 3 <211> 20 <212> PRT <213> Mus musculus <400> 3 Pro Ala Cys Lys Asn Val Asp Ala Asp Glu Asp Glu Asp Tyr Pro Asn Gly Phe Leu <210> 4 <211> 24

<212> DNA

<213> artificial sequence

<220>



PCT/IB03/01044

6

<223> primer

<400> 4 gtggcaagct acgagaacca gggt

24

<210> 5

<211> 24

<212> DNA

<213> artificial sequence

<220>

<223> primer

<400> 5 gacgaaggag caaaggtgga agga

24

<210> 6

<211> 22

<212> DNA

<213> artificial sequence

<220>

<223> primer

<400> 6 cccagaggca aaccctctga ag

22

<210> 7

<211> 24

<212> DNA

<213> artificial sequence

<220>





7

<223> primer

<400> 7 tcgaattcgc caatgacaag acgc

24

<210> 8

<211> 21

<212> DNA

<213> artificial sequence

<220>

<223> primer

<400> 8 ggagacttag atgtctgacc g

21

<210> 9

<211> 21

<212> DNA

<213> artificial sequence

<220>

<223> primer

<400> 9 gacagaccag cagggacagt g

21





inter al Application No PCT/IB 03/01044

A. CLASSIFICATION OF SUBJECT MATTER
IPC 7 C12N15/12 C12N15/63 C07K14/47
G01N33/68

CO7K14/47 A01K67/027 C07K16/18

According to international Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

EMBL, SEQUENCE SEARCH, EPO-Internal, WPI Data, PAJ, BIOSIS, MEDLINE, CHEM ABS Dat

Category •	Citation of document, with indication, where appropriate, of the	relevant passages	Relevant to claim No.
X	SOMMERS CONNIE L ET AL: "Knock mutation of the distal four tyr linker for activation of T cell murine T cell development." JOURNAL OF EXPERIMENTAL MEDICIN vol. 194, no. 2, 16 July 2001 (2001-07-16), page XP002207177 ISSN: 0022-1007 cited in the application the whole document	osines of s blocks	1,5-8, 11-13, 15,22
X Furth	er documents are listed in the continuation of box C.	X Patent family members are listed	in annex.
"A" docume conside "E" earlier of fling d' "L" docume which in citation "O" docume other n' "P" docume	nt which may throw doubts on priority ctalm(s) or is cited to establish the publication date of another or other special reason (as specified) intreferring to an oral disclosure, use, exhibition or	"T' tater document published after the inte or priority date and not in conflict with cited to understand the principle or th invention "X" document of particular relevance; the cannot be considered novel or carno involve an inventive step when the document of particular relevance; the cannot be considered to involve an in document is combined with one or my ments, such combination being obvio in the art. "8" document member of the same patent	the application but econy underlying the considered to coment is taken alone laimed invention ventive step when the one other such docu— us to a person skilled
	actual completion of the international search July 2003	Date of mailing of the international set	arch report
	nalling address of the ISA European Patent Cifice, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Authorized officer Devijver, K	



INTERNATIONAL SEARCH REPORT

PCT/IB 03/01044

	PC1/1B 03/01044
	Relevant to claim No.
Citation of document, with indication, where appropriate, or the resevant passages	ndevall to claim two.
AGUADO ENRIQUE ET AL: "Induction of T helper type 2 immunity by a point mutation in the LAT adaptor." SCIENCE (WASHINGTON D C), vol. 296, no. 5575, 2002, pages 2036-2040, XP002207182 14 June, 2002 ISSN: 0036-8075 the whole document -& DATABASE EMBL 'Online! 2 May 2002 (2002-05-02) MALISSEN M: "Mus musculus LAT gene for linker protein, exons 1-12." Database accession no. AJ438435 XP002207184	1-34
SOMMERS CONNIE L ET AL: "A LAT mutation that inhibits T cell development yet induces lymphoproliferation." SCIENCE (WASHINGTON D C), vol. 296, no. 5575, 2002, pages 2040-2043, XP002249670 14 June, 2002 ISSN: 0036-8075 the whole document	1-34
SAMELSON L E ET AL: "STUDIES ON THE ADAPTER MOLECULE LAT" COLD SPRING HARBOR SYMPOSIA ON QUANTITATIVE BIOLOGY, BIOLOGICAL LABORATORY, COLD SPRING HARBOR, NY, US, no. 64, 1999, pages 259-263, XP001056468 ISSN: 0091-7451 cited in the application page 261	1-34
LIN JOSEPH ET AL: "Identification of the minimal tyrosine residues required for linker for activation of T cell function." JOURNAL OF BIOLOGICAL CHEMISTRY, vol. 276, no. 31, 3 August 2001 (2001–08–03), pages 29588-29595, XP002207178 ISSN: 0021-9258 cited in the application the whole document	1-34
	helper type 2 immunity by a point mutation in the LAT adaptor." SCIENCE (WASHINGTON D C), vol. 296, no. 5575, 2002, pages 2036-2040, XP002207182 14 June, 2002 ISSN: 0036-8075 the whole document -& DATABASE EMBL 'Online! 2 May 2002 (2002-05-02) MALISSEN M: "Mus musculus LAT gene for linker protein, exons 1-12." Database accession no. AJ438435 XP002207184 SOMMERS CONNIE L ET AL: "A LAT mutation that inhibits T cell development yet induces lymphoproliferation." SCIENCE (WASHINGTON D C), vol. 296, no. 5575, 2002, pages 2040-2043, XP002249670 14 June, 2002 ISSN: 0036-8075 the whole document SAMELSON L E ET AL: "STUDIES ON THE ADAPTER MOLECULE LAT" COLD SPRING HARBOR SYMPOSIA ON QUANTITATIVE BIOLOGY, BIOLOGICAL LABORATORY, COLD SPRING HARBOR, NY, US, no. 64, 1999, pages 259-263, XP001056468 ISSN: 0091-7451 cited in the application page 261 LIN JOSEPH ET AL: "Identification of the minimal tyrosine residues required for linker for activation of T cell function." JOURNAL OF BIOLOGICAL CHEMISTRY, vol. 276, no. 31, 3 August 2001 (2001-08-03), pages 29588-29595, XP002207178 ISSN: 0021-9258 cited in the application the whole document



INTERNATIONAL SEARCH REPORT

Inte inal Application No PCT/IB 03/01044

Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT tegory * Citation of document, with indication, where appropriate, of the relevant passages Relevant to dalim No.					
ZHANG WEIGUO ET AL: "Association of Grb2, Gads, and phospholipase C-gammal with phosphorylated LAT tyrosine residues. Effect of LAT tyrosine mutations on T cell antigen receptor-mediated signaling." JOURNAL OF BIOLOGICAL CHEMISTRY, vol. 275, no. 30, 28 July 2000 (2000-07-28), pages 23355-23361, XP002207179 ISSN: 0021-9258 cited in the application the whole document	1-34				
WO 99 32627 A (SAMELSON LAWRENCE E ;US HEALTH (US); ZHANG WEIGUO (US)) 1 July 1999 (1999-07-01) * SEQ ID NO:3; SEQ ID NO:5 * claims 1-25	1-34				
ZHANG WEIGUO ET AL: "Essential role of LAT in T cell development." IMMUNITY, vol. 10, no. 3, March 1999 (1999-03), pages 323-332, XP002207180 ISSN: 1074-7613					
SAITOH SHINICHIROH ET AL: "LAT is essential for FcepsilonRI-mediated mast cell activation." IMMUNITY, vol. 12, no. 5, May 2000 (2000-05), pages 525-535, XP002207181 ISSN: 1074-7613					

INTERNATIONAL SEARCH REPORT

PCT/IB 03/01044

Box I	Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This inte	emational Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. X	Claims Nos.: because they relate to subject matter not required to be exarched by this Authority, namely:
	Although claim 32 is directed to a diagnostic method practised on the human/animal body, the search has been carried out and based on the alleged effects of the compound/composition.
2. X	Claims Nos.: 21 because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically: See FURTHER INFORMATION sheet PCT/ISA/210
	see Further information sheet PC1/15A/210
3.	Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II	Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This inte	emational Searching Authority found multiple inventions in this international application, as follows:
1.	As all required additional search fees were timely paid by the applicant, this international Search Report covers all searchable claims.
2.	As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3.	As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:
	·
4.	No required additional search fees were timely paid by the applicant. Consequently, this international Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark	t on Protest The additional search fees were accompanied by the applicant's protest.
	No protest accompanied the payment of additional search fees.

International Application No. PCT/IB 03 01044

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

Continuation of Box I.2

Claims Nos.: 21

Claim 21 refers to a compound identified using a screening method according to any of claims 16 to 20 without giving a true technical characterization. Moreover, no such compounds are defined in the application. In consequence, the scope of said claim is ambiguous and vague, and its subject-matter is not sufficiently disclosed and supported (Art. 5 and 6 PCT). No meaningful search can be carried out for such a purely speculative claim whose wording is, in fact, a mere recitation of the result to be achieved.

The applicant's attention is drawn to the fact that claims, or parts of claims, relating to inventions in respect of which no international search report has been established need not be the subject of an international preliminary examination (Rule 66.1(e) PCT). The applicant is advised that the EPO policy when acting as an International Preliminary Examining Authority is normally not to carry out a preliminary examination on matter which has not been searched. This is the case irrespective of whether or not the claims are amended following receipt of the search report or during any Chapter II procedure.





Inter Ial Application No PCT/IB 03/01044

Patent document died in search report		Publication date		Patent family member(s)	Publication date
WO 9932627	A	01-07-1999	AU AU CA EP WO	750543 B2 2204799 A 2316769 A1 1141281 A2 9932627 A2	18-07-2002 12-07-1999 01-07-1999 10-10-2001 01-07-1999